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NEWS 9 APR 30 CHEMCATS enhanced with 1.2 million new records
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NEWS 13 MAY 08 CA/Capplus Indian patent publication number format defined
NEWS 14 MAY 14 RDISCLOSURE on STN Easy enhanced with new search and display fields
NEWS 15 MAY 21 BIOSIS reloaded and enhanced with archival data
NEWS 16 MAY 21 TOXCENTER enhanced with BIOSIS reload
NEWS 17 MAY 21 CA/Capplus enhanced with additional kind codes for German patents
NEWS 18 MAY 22 CA/Capplus enhanced with IPC reclassification in Japanese patents
NEWS 19 JUN 27 CA/Capplus enhanced with pre-1967 CAS Registry Numbers
NEWS 20 JUN 29 STN Viewer now available
NEWS 21 JUN 29 STN Express, Version 8.2, now available
NEWS 22 JUL 02 LEMBASE coverage updated
NEWS 23 JUL 02 LMEDLINE coverage updated
NEWS 24 JUL 02 SCISEARCH enhanced with complete author names
NEWS 25 JUL 02 CHEMCATS accession numbers revised
NEWS 26 JUL 02 CA/Capplus enhanced with utility model patents from China
NEWS 27 JUL 16 Capplus enhanced with French and German abstracts
NEWS 28 JUL 18 CA/Capplus patent coverage enhanced
NEWS 29 JUL 26 USPATFULL/USPAT2 enhanced with IPC reclassification
NEWS 30 JUL 30 USGENE now available on STN

NEWS EXPRESS 29 JUNE 2007: CURRENT WINDOWS VERSION IS V8.2,
CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
AND CURRENT DISCOVER FILE IS DATED 05 JULY 2007.

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SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

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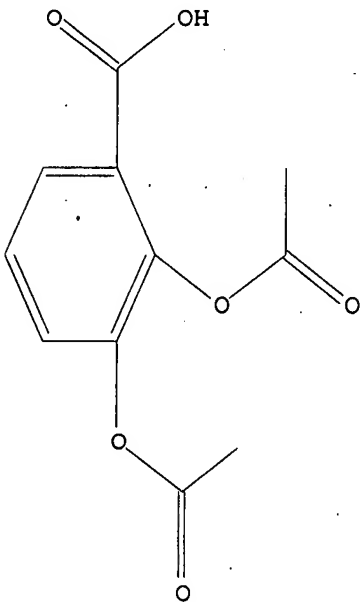
Uploading C:\Program Files\Stnexp\Queries\2,3-diacetoxy2.str

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s l1 fam sam

SAMPLE SEARCH INITIATED 14:40:51 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 10 TO ITERATE

100.0% PROCESSED 10 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 11 TO 389
PROJECTED ANSWERS: 0 TO 0

L2 0 SEA FAM SAM L1

=> s l1 fam full

FULL SEARCH INITIATED 14:40:56 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 131 TO ITERATE

100.0% PROCESSED 131 ITERATIONS 2 ANSWERS
SEARCH TIME: 00.00.01

L3 2 SEA FAM FUL L1

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COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	67.70	67.91

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L4 53 L3

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L5 3 ISCHMI?

=> s ischemi?

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L6 89656 ISCHEMI?

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L7 1 L4 AND L6

=> d ti au abs so py

L7 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN
TI Use of 2,3-alkylcarbonyloxybenzoic acids, derivatives and analogues
therefrom in the treatment of tissue and cellular dysfunction, damage and
injury in mammals
IN Stec, Karen; Rubinstein, Israel; Eiznhamer, David; Xu, Ze-qu; Flavin,
Michael
AB A method for the treatment of cellular and tissue damage is disclosed.
The inventive method comprises the use of 2,3-alkylcarbonyloxybenzoic acid
and salts thereof for the prevention and treatment of dysfunction, damage,
and/or injuries to organs, tissues and/or cells in human or animal
subjects caused by diseases, infections and conditions such as pneumonia,
coronavirus, multiple transfusions, trauma, ischemic-reperfusion
dysfunctions, stroke, drug overdose, and severe acute respiratory
syndrome. The 2;3-alkylcarbonyloxybenzoic acid may be used alone or in
combination with other therapeutic agents such as antibiotics. The acid
may be administered in any practical delivery form, and in free acid or
buffered form.
SO U.S. Pat. Appl. Publ., 6 pp.
CODEN: USXXCO
PY 2004
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=> d ti au abs so py 1-10 14

L4 ANSWER 1 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Treatment of VR1-antagonist-induced increase in body temperature with an
antipyretic agent
IN Bannon, Anthony W.; Beck, Klaus D.; Treanor, James J. S.
AB The invention relates to a method of reducing a VR1-antagonist-induced
increase in body temperature in a mammal in need thereof, comprising the step
of
administering an antipyretic agent to the mammal and the like. TRPV1
antagonist treatment of rats resulted in hyperthermia which was reversed
by acetaminophen administration.
SO PCT Int. Appl., 151pp.
CODEN: PIXXD2
PY 2006
2006

L4 ANSWER 2 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Novel nanoparticulate nimesulide compositions
IN Bosch, H. William; Wertz, Christian F.
AB The present invention provides nanoparticulate nimesulide compns. The
compns. preferably comprise nimesulide and at least one surface stabilizer
adsorbed on or associated with the surface of the nimesulide particles. The
nanoparticulate nimesulide particles preferably have an effective average
particle size of less than about 2000 nm. The invention also provides
methods of making and using nanoparticulate nimesulide compns. An aqueous
solution of 1% (weight/weight) Plasdone S-630 was combined with 4.25 g of
nimesulide (5% weight/weight) and stirred for 1 h at 4200 rpm with chilled
water
(10°) recirculated through the milling chamber. The process
yielded a colloidal dispersion of nimesulide with a mean particle size of
150 nm, a D50 of 124 nm, a D90 of 256 nm, and a D95 of 293 nm.
SO PCT Int. Appl., 87 pp.
CODEN: PIXXD2
PY 2005
2005
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2006

L4 ANSWER 3 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Inductive QSAR descriptors. Distinguishing compounds with antibacterial activity by artificial neural networks
 AU Cherkasov, Artem
 AB On the basis of the previous models of inductive and steric effects, 'inductive' electronegativity and mol. capacitance, a range of new 'inductive' QSAR descriptors has been derived. These mol. parameters are easily accessible from electronegativities and covalent radii of the constituent atoms and interat. distances and can reflect a variety of aspects of intra- and intermol. interactions. Using 34 'inductive' QSAR descriptors alone we have been able to achieve 93% correct separation of compds. with- and without antibacterial activity (in the set of 657). The elaborated QSAR model based on the Artificial Neural Networks approach has been extensively validated and has confidently assigned antibacterial character to a number of trial antibiotics from the literature.
 SO International Journal of Molecular Sciences (2005), 6(1-2), 63-86
 CODEN: IJMCFK; ISSN: 1422-0067
 URL: <http://www.mdpi.org/ijms/papers/i6010063.pdf>
 PY 2005

L4 ANSWER 4 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Dispersible formulations containing anti-inflammatory agents and other active ingredients for infusion
 IN Britten, Nancy Jean; Waldron, Niki Ann; Watts, Jeffrey L.; Hallberg, John Walter; Burns, John W.
 AB A method is provided for treatment and/or prevention of an inflammatory condition in a fluid-containing organ having a natural exterior orifice, such as the udder of a milk-producing animal or an ear of a subject. The invention also relates to a dispersible pharmaceutical composition suitable for infusion into the organ according to the method of the invention, and a process for preparing such a composition. For example, a suspension to be administered by intrammary infusion was prepared containing parecoxib 100 mg/mL, Labrafil M-1944CS 50 mg/mL, microcryst. wax 70 mg/mL,, and cottonseed oil q.s.
 SO U.S. Pat. Appl. Publ., 22 pp., Cont.-in-part of U.S. Ser. No. 803,146.
 CODEN: USXXCO
 PY 2005
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L4 ANSWER 5 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Dispersible pharmaceutical composition for treatment of mastitis and otic disorders
 IN Britten, Nancy J.; Burns, John W.; Hallberg, John W.; Waldron, Niki A.; Watts, Jeffrey L.
 AB A method is provided for treatment of an infective condition in a fluid-containing organ having a natural exterior orifice, such as the udder of a milk producing animal or an ear. The method comprises administering an antibacterial agent to the organ via the exterior orifice and administering in combination therapy with the antibacterial agent a second agent that is an anti-inflammatory agent, an analgesic and/or an antipyretic. The antibacterial agent and, optionally, the second agent, are administered as a pharmaceutical composition further comprising a vehicle that comprises an amphipathic oil that is water dispersible and ethanol

insol., microcryst. wax and a pharmaceutically acceptable non-aqueous carrier. Also provided is such a composition comprising the antibacterial agent and the second agent. The composition is readily dispersible in the fluid of the fluid-containing organ. A suspension to be administered by intramammary infusion was contained ceftiofur hydrochloride (micronized) 12.5 mg/mL, Labrafil M-1944CS 50 mg/mL, microcryst. wax 100 mg/mL, cottonseed oil q.s.

SO PCT Int. Appl., 58 pp.

CODEN: PIXXD2

PY

2004
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L4 ANSWER 6 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

TI Dispersible formulations of an anti-inflammatory agent

IN Britten, Nancy J.; Burns, John W.; Hallberg, John W.; Waldron, Niki A.; Watts, Jeffrey L.

AB A method is provided for treatment of an inflammatory condition in a fluid-containing organ having a natural exterior orifice, such as the udder of a milk producing animal or an ear. The method comprises administering, to the organ via the exterior orifice, a pharmaceutical composition comprising an anti-inflammatory agent and a vehicle that comprises an amphipathic oil that is water dispersible and ethanol insol., microcryst. wax and a pharmaceutically acceptable non-aqueous carrier. Also provided is such a composition comprising the anti-inflammatory agent. The composition is readily dispersible in the fluid of the fluid-containing organ. Thus, a suspension to be administered by intramammary infusion comprised parecoxib 100, Labrafil M-1944CS 50, and microcryst. wax 70 mg/mL, and cottonseed oil qs.

SO PCT Int. Appl., 45 pp.

CODEN: PIXXD2

PY

2004
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L4 ANSWER 7 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

TI Use of 2,3-alkylcarbonyloxybenzoic acids in the treatment of anthrax

IN Stec, Karen J.

AB A method for treating inhalation anthrax is disclosed. The inventive method comprises the use of 2,3-alkylcarbonyloxybenzoic acid and salts thereof in the prevention and treatment of lung damage caused by Bacillus anthracis and toxins produced by the bacterium. The 2,3-alkylcarbonyloxybenzoic acid may be used alone or in combination with other therapeutic agents such as antibiotics.

SO U.S. Pat. Appl. Publ., 3 pp.

CODEN: USXXCO

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2004
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L4 ANSWER 8 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Effective attenuation of endotoxin-induced acute lung injury by
 2,3-diacetyloxybenzoic acid in two independent animal models
 AU Eiznhamer, David A.; Flavin, Michael T.; Jesmok, Gary J.; Borgia, Julian
 F.; Nelson, Deanna J.; Burhop, Kenneth E.; Xu, Ze-Qi
 AB The pathol. of acute lung injury (ALI) is often modeled in animal studies
 by the administration of lipopolysaccharide (LPS), which results in an
 endotoxemia with sequelae similar to that seen in acute respiratory
 distress syndrome (ARDS). Here we report the results of two studies
 designed to examine the efficacy of a novel agent, 2,3-diacetyloxybenzoic
 acid (2,3-DABA), in the treatment of LPS-induced ALI. In two sep. animal
 models, 2,3-DABA was effective in significantly reducing lung
 microvascular permeability, a condition commonly seen in ARDS, which
 results in pulmonary edema and respiratory insufficiency. In each model,
 it is demonstrated that the mechanism by which 2,3-DABA exerts this effect
 occurs subsequent to the recruitment of neutrophils to the site of
 inflammation. Lung permeability was significantly decreased in both
 models by treatment with 2,3-DABA, suggesting that this agent, either
 alone or in combination therapy, may be useful in the treatment of ALI
 associated with ARDS.
 SO Pulmonary Pharmacology & Therapeutics (2004), 17(2), 105-110
 CODEN: PPTHFJ; ISSN: 1094-5539
 PY 2004

L4 ANSWER 9 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Use of 2,3-alkylcarbonyloxybenzoic acids, derivatives and analogues
 therefrom in the treatment of tissue and cellular dysfunction, damage and
 injury in mammals
 IN Stec, Karen; Rubinstein, Israel; Eiznhamer, David; Xu, Ze-qu; Flavin,
 Michael
 AB A method for the treatment of cellular and tissue damage is disclosed.
 The inventive method comprises the use of 2,3-alkylcarbonyloxybenzoic acid
 and salts thereof for the prevention and treatment of dysfunction, damage,
 and/or injuries to organs, tissues and/or cells in human or animal
 subjects caused by diseases, infections and conditions such as pneumonia,
 coronavirus, multiple transfusions, trauma, ischemic-reperfusion
 dysfunctions, stroke, drug overdose, and severe acute respiratory
 syndrome. The 2,3-alkylcarbonyloxybenzoic acid may be used alone or in
 combination with other therapeutic agents such as antibiotics. The acid
 may be administered in any practical delivery form, and in free acid or
 buffered form.
 SO U.S. Pat. Appl. Publ., 6 pp.
 CODEN: USXXCO
 PY 2004
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 2004
 2005

L4 ANSWER 10 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
 TI Discrimination and selection of new potential antibacterial compounds
 using simple topological descriptors
 AU Murcia-Soler, Miguel; Perez-Gimenez, Facundo; Garcia-March, Francisco J.;
 Salabert-Salvador, M. Teresa; Diaz-Villanueva, Wladimiro;
 Medina-Casamayor, Piedad
 AB The aim of the work was to discriminate between antibacterial and
 non-antibacterial drugs by topol. methods and to select new potential
 antibacterial agents from among new structures. The method used for
 antibacterial activity selection was a linear discriminant anal. (LDA).
 It is possible to obtain a QSAR interpretation of the information
 contained in the discriminant function. We make use of the pharmacol.
 distribution diagrams (PDDs) as a visualizing technique for the
 identification and selection of new antibacterial agents.
 SO Journal of Molecular Graphics & Modelling (2003), 21(5), 375-390
 CODEN: JMGMFJ; ISSN: 1093-3263

PY 2003

=> d ti au abs so py 11-20 14

- L4 ANSWER 11 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Structure-Based Classification of Antibacterial Activity
AU Cronin, Mark T. D.; Aptula, Aynur O.; Dearden, John C.; Duffy, Judith C.;
Netzeva, Tatiana I.; Patel, Hiren; Rowe, Philip H.; Schultz, T. Wayne;
Worth, Andrew P.; Voutzoulidis, Konstantinos; Schueuermann, Gerrit
AB The aim of this study was to develop a simple quant. structure-activity
relation (QSAR) for the classification and prediction of antibacterial
activity, to enable in silico screening. To this end a database of 661
compds., classified according to whether they had antibacterial activity,
and for which a total of 167 physicochem. and structural descriptors were
calculated, was analyzed. To identify descriptors that allowed separation of
the two classes (i.e. those compds. with and without antibacterial activity),
anal. of variance was utilized and models were developed using linear
discriminant and binary logistic regression analyses. Model predictivity
was assessed and validated by the random removal of 30% of the compds. to
form a test set, for which predictions were made from the model. The
results of the analyses indicated that six descriptors, accounting for
hydrophobicity and inter- and intramol. hydrogen bonding, provided
excellent separation of the data. Logistic regression anal. was shown to model
the data slightly more accurately than discriminant anal.
SO Journal of Chemical Information and Computer Sciences (2002), 42(4),
869-878
CODEN: JCISD8; ISSN: 0095-2338
PY 2002
- L4 ANSWER 12 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Synthesis and biological properties of 3-(dihydroxybenzoyloxy)methyl- and
3-(diacetoxybenzoyloxy)-methyl-7 α -chlorocephalosporanate sulfones
AU Grigan, N.; Veinberg, G.; Shestakova, I.; Kanepe, I.; Lukevics, E.
AB The synthesis of tert-Bu esters of 3-(2-hydroxybenzoyloxy)methyl-,
3-(dihydroxybenzoyloxy)methyl-, and 3-(diacetoxybenzoyloxy)methyl-7 α -
chlorocephalosporanic acid sulfones by reaction of tert-Bu ester of
3-bromomethyl-7 α -chlorocephalosporanic acid sulfone with salts of
hydroxy- and acetoxy-substituted benzoic acids is described. The
elastase-inhibiting properties of the compds. obtained and also their in
vitro cytotoxic activity were investigated.
SO Chemistry of Heterocyclic Compounds (New York, NY, United
States) (Translation of Khimiya Geterotsiklicheskikh Soedinenii) (2000),
36(10), 1232-1236
CODEN: CHCCAL; ISSN: 0009-3122
PY 2000
- L4 ANSWER 13 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Synthesis, activity and formulations of pharmaceutical compounds for
treatment of oxidative stress and/or endothelial dysfunction
IN Del Soldato, Piero
AB Compds. or their salts of general formula (I): A-B-N(O)_s wherein: s is an
integer equal to 1 or 2; A = R-Tl-, wherein R is the drug radical and Tl =
(CO)t or (X)t', wherein X = O, S, NRlc, Rlc is H or a linear or branched
alkyl or a free valence, t and t' are integers and equal to zero or 1,
with the proviso that t = 1 when t' = 0; t = 0 when t' = 1; B = -TB -X₂-O-
wherein TB = (CO) when t = 0, TB = X when t' = 0, X being as above
defined; X₂, bivalent radical, is such that the precursor drug of A and
the precursor of B meet resp. the pharmacol. tests described in the
description. Synthesis, activity and formulations of pharmaceutical
compds. for treatment of oxidative stress and/or endothelial dysfunction
are disclosed. The precursors are such as to meet the pharmacol. test
reported in the description.

SO PCT Int. Appl., 94 pp.

CODEN: PIXXD2

PY 2001

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L4 ANSWER 14 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

TI Synthesis, activity and formulations of pharmaceutical compounds for treatment of oxidative stress and/or endothelial dysfunction

IN Del Soldato, Piero

AB Synthesis, activity and formulations of pharmaceutical compds. for treatment of oxidative stress and/or endothelial dysfunction are disclosed. The precursors are such as to meet the pharmacol. test reported in the description.

SO PCT Int. Appl., 140 pp.

CODEN: PIXXD2

PY 2000

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L4 ANSWER 15 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

TI Synthesis, activity and formulations of pharmaceutical compounds for treatment of oxidative stress and/or endothelial dysfunction

IN Del Soldato, Piero

AB Compds. A-B-C-N(O)s and A-C1[N(O)s]-B1 or their salts [s is an integer 1

or 2, preferably $s = 2$; A is the radical of a drug and is such as to meet the pharmacol. tests reported in the description; C and Cl are two bivalent radicals; the precursors of the radicals B and Bl are such as to meet the pharmacol. test reported in the description] were prepared for use as pharmaceuticals. Thus, (S,S)-N-acetyl-S-(6-methoxy- α -methyl-2-naphthalenylacetyl)cysteine 4-nitroxybutyl ester was prepared (NCX 2101) from naproxene and N-acetylcysteine in the first of 28 synthetic examples given. Pharmacol. test examples and tabular data are also given.

SO PCT Int. Appl., 159 pp.

CODEN: PIXXD2

PY 2000

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L4 ANSWER 16 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

TI Synthesis and in vitro antibacterial activity of catechol-spiramycin conjugates

AU Poras, Herve; Kunesch, Gerhard; Barriere, Jean-Claude; Berthaud, Nadine; Andreumont, Antoine

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The first synthesis of siderophore conjugates of two macrolide antibiotics, spiramycin (I) and neospiramycin (II), which are unable to penetrate the outer membrane of Gram-neg. bacteria are described. These novel conjugates were prepared by regioselective acylation of a hydroxyl function of I and II with a dihydroxybenzoic Fe(III) complexing ligand linked via a carboxyl group containing spacer to the macrolide antibiotics. The preliminary biol. evaluation of these novel conjugates under standard and iron depleted conditions has shown that their antibacterial activity was comparable to that of I and II.

SO Journal of Antibiotics (1998), 51(8), 786-794

CODEN: JANTAJ; ISSN: 0021-8820

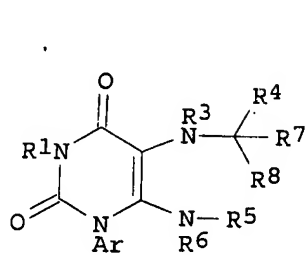
PY 1998

L4 ANSWER 17 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN

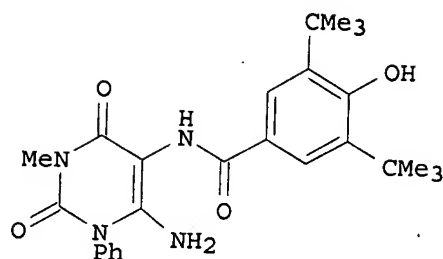
TI Preparation of 1-arylpyrimidine derivatives as antiallergics.

IN Isobe, Yoshiaki; Katagiri, Toshimasa; Umezawa, Junko; Goto, Yuso; Sasaki, Masashi; Watanabe, Nobuo; Sato, Hideharu; Obara, Fumihiko

GI



I



II

- AB The invention relates to 1-arylpyrimidine derivs. I [R1 = H, alkyl, or aralkyl; Ar = 1-naphthyl or (un)substituted Ph; R4 = substituted Ph, substituted styryl, 1-methylcyclohexyl, 4-methylcyclohexyl, 4-oxo-4H-pyran-2-yl, or 2-oxo-2H-pyran-5-yl; R5, R6 = H or alkyl; R3 = H and R7R8 = oxo; or R3R7 = bond and R5R8 = bond], or pharmaceutically acceptable salts thereof, and their use as agents for treating allergic diseases. For example, reaction of 5,6-diamino-3-methyl-1-phenyluracil with 4-hydroxy-3,5-di-tert-butylbenzoyl chloride [prepns. given] in CHCl3 containing pyridine gave 79% title compound II. In tests for inhibition of picryl chloride-induced type IV allergy in mice and PCA in rats, I were comparable to the pos. stds. prednisolone and tranilast. Toxic effects were not observed in rats given I at oral dosages of 1000 mg/kg/day for 2 wk.
- SO Can. Pat. Appl., 104 pp.
CODEN: CPXXEB
- PY 1996
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- L4 ANSWER 18 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
- TI Use of 2,3 alkylcarbonyloxybenzoic acid in treating adult respiratory distress syndrome
- IN Flavin, Michael T.; Nelson, Deanna J.; Borgia, Deceased Julian F.; Jesmok, Gary
- AB Methods for treating adult respiratory distress syndrome (ARDS) which involves the administration of C2-18 2,3-alkylcarbonyloxybenzoic acids and salts are described. The therapeutic efficacy of 2,3-diacetoxybenzoic acid in combination with ibuprofen eas demonstrated in an ARDS sheep model.
- SO U.S., 6 pp.
CODEN: USXXAM
- PY 1996
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- L4 ANSWER 19 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
- TI Method and use of agents to inhibit protein polymerization, methods of identifying these agents, and use of the agents as antithrombotics and for the treatment of Alzheimer's disease

IN Bjornsson, Thorir D.
AB A method of inhibiting polymerization of target proteins by administration of compds. capable of inhibiting aggregation and subsequent transglutaminase-induced crosslinking of adjacent peptides of the target proteins is provided. These compds. are useful as antithrombotic agents and in the treatment of Alzheimer's disease. A method of screening and identifying compds. capable of inhibiting aggregation and subsequent transglutaminase-induced crosslinking of amyloid β -peptide is also provided.
SO PCT Int. Appl., 18 pp.
CODEN: PIXXD2
PY 1995

L4 ANSWER 20 OF 53 CAPLUS COPYRIGHT 2007 ACS on STN
TI Growth promotion of synthetic catecholate derivatives on Gram-negative bacteria
AU Reissbrodt, Rolf; Heinisch, Lothar; Mollmann, Ute; Rabsch, Wolfgang; Ulbricht, Hermann
AB Derivs. of benzoic acid, glyoxylic acid benzhydrazone, oxanilic acid and N-dihydroxybenzylidene-2,4,6-trimethylaminobenzene were investigated as catecholic iron chelators under iron-depleted conditions. Some of the compds. showed strong pos. reactions in the universal chemical siderophore assay (CAS): 3,4-dihydroxybenzoic acid, glyoxylic acid 2,3-dihydroxybenzhydrazone, N-3,4-dihydroxybenzylidene-2,4,6-trimethylaminobenzene. In particular these compds. also enabled removal of iron from iron-saturated transferrin. Using various siderophore indicator strains (Enterobacteriaceae, Pseudomonas aeruginosa and Aeromonas hydrophila mutants) in bioassays the following growth promotion could be detected: vicinal substituents (e.g. 2,3- or 3,4-) were essential, the carboxyamido group seen in benzoic acids and glyoxylic acid benzhydrazones contributed to a pos. reaction as well as the azomethine group (in N-3,4-dihydroxybenzylidene-2,4,6-trimethylaminobenzene). 2,3-Dihydroxybenzoic acid and the 2,3-diacetoxy substitute preferably promoted growth of Enterobacteriaceae mutants. In contrast, the 3,4-positioned compds. preferably promoted growth of P. aeruginosa mutants and A. hydrophila SB 22. Glyoxylic acid di(methoxycarbonyloxy)-benzhydrazones (2,3- and 3,4- positioned) including the 2,3-dihydroxy compound preferably enabled growth of the non-fermenters. N-3,4-dihydroxybenzylidene-2,4,6-trimethylaminobenzene supplied all mutants of Salmonella, Escherichia coli, Klebsiella, Morganella, P. aeruginosa and A. hydrophila with iron. Transport of glyoxylic acid 2,3-dihydroxybenzhydrazone depended on tonB, and required the involvement of the iron-regulated outer membrane proteins (IROMPs) FepA, Cir and Fiu.
SO BioMetals (1993), 6(3), 155-62
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EAST Search History

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	3	wo "2004032825"	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2007/08/03 14:33
L2	5	2,3-diacetoxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2007/08/03 14:48
L3	143	del-soldato-piero.in.	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2007/08/03 14:48
S1	5	2,3-diacetoxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2007/08/03 14:27
S2	10	2,3-alkylcarbonyloxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 14:31
S3	74	dipyrrocetyl	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/11/20 13:27
S4	7	2,3-diacetoxybenzoic adj acid or 2,3-DABA or "2,3" adj DABA	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2006/12/11 14:27
S5	8364	ischemia near reperfusion	US-PGPUB; USPAT; EPO; JPO; DERWENT	NEAR	ON	2006/12/11 14:31
S6	1	S4 and S5	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 14:31
S7	10	2,3-alkylcarbonyloxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 14:32
S8	2	S7 and S5	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 14:32

EAST Search History

S9	74	dipyrocetyl	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 15:07
S10	1	S9 and S5	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 14:32
S11	663	alteplase	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 15:07
S12	0	S9 and S11	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 15:07
S13	1	S7 and S11	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 15:07
S14	1	S4 and S11	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 15:07

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S2	10	2,3-alkylcarbonyloxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 14:31
S3	74	dipyroceryl	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/11/20 13:27
S4	7	2,3-diacetoxybenzoic adj acid or 2,3-DABA or "2,3" adj DABA	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2006/12/11 14:27
S5	8364	ischemia near reperfusion	US-PGPUB; USPAT; EPO; JPO; DERWENT	NEAR	ON	2006/12/11 14:31
S6	1	S4 and S5	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 14:31
S7	10	2,3-alkylcarbonyloxybenzoic adj acid	US-PGPUB; USPAT; EPO; JPO; DERWENT	ADJ	ON	2006/12/11 14:32
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S14	1	S4 and S11	US-PGPUB; USPAT; EPO; JPO; DERWENT	AND	ON	2006/12/11 15:07